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(54) Title: SUBTILISIN MUTANTS

(57) Abstract

Novel carbonyl hydro-DNA sequences of naturallynon-human carbonyl hydroant carbonyl hydrolases, in recombinant carbonyl hydrolase to generate the substituid residues in the amino acid sequence of a precursor carbonyl hydrolase. Such mutant porperties which are different from those of the precursor hydrolase and are especially tions. The substituted amino acid residues correspond to position +123 and/or +274 in Bacillus amyloliquefaciens subtilisin.

1 GCG CAA TCA GTG CCA TGG GGC ATC TCG CGA GTT CAA GCT CCT GCT GCT CAC AAC CGC GGC 1 Ala Gln Ser Val Pro Trp Gly Ile Ser Arg Val Gin Ala Pro Ala Ala His Asn Arg Gly 61 TTA ACA GGC AGC GGC GTT AGA GTT GCT GTT TTA GAT ACA GGC ATC AGC ACA --- CAC CCA 21 Leu Thr Gly Ser Gly Val Arg Val Ala Val Leu Asp Thr Gly 1le Ser Thr - His Pro lase mutants derived from the
121 GAT CTT AAT ATT AGA GGC GGC GCG AGC TTC GTT CCC GGC GAA --- CCG TCG ACA CAA GAT
DNA sequences of naturally lases are disclosed. The mut- 241 GTT TTA GGC GTT GCT CCT TCG GCC GAA TTA TAC GCT GTT AAA GTT-TTA GGC GCT AGC GGC ant carbonyl hydrolases in 81 Val Leu Gly Val Ala Pro Ser Ala Glu Leu Tyr Ala Val Lys Vali Leu Gly Ala Ser Gly general, are obtained by in 301 AGC GGC AGC TAC AGC TCT ATC GCT CAA GGC CTC GAG TGG GCT GGC AAC AAC GGT AGT CAC vitro modification of a precursor DNA sequence encod- 361 GTT GCT AGC TTA AGC CTC GGG AGC CCT AGC CCT AGC GCT ACA TTA GAA CAA GCT GTT AAC ing the naturally-occurring or 121 Val Ala Ser Leu Ser Leu Gly Ser Pro Ser Pro Ser Ala Thr Leu Glu Gln Ala Val Asn lase to generate the substitu421 AGC GCT ACA TCT AGA GGC GTT TTA GTT GCT GCG AGC GGC AAC AGC GGC GGT GGA TCG
tion of one or more amino ac141 Ser Ala Thr Ser Arg Gly Val Leu Val Val Ala Ala Ser Gly Asn Ser Gly Ala Gly Ser 481 --- --- ATC AGC CTA CCC TGC TAG ATA CGC TAA TGC CAT GGC TGT TGG CGC ACA 161 - - - - - Ile Ser tyr Pro Ala Arg Tyr Ala Asn Ala Met Ala Val Gly Ala Thr onyl hydrolase. Such mutant 541 gat caa aac aac aga gca agc ttc agt caa tac ggc gct ggc tta gat atc gtg gcg carbonyl hydrolases have 181 asp gin asn asn asn arg ala Ser Phe Ser gin tyr gly ala gly Leu asp lie val ala 601 CCT GGC GTT AAC GTT CAA AGC ACA TAC CCT GGC AGC ACA TAC GCC AGC TTG AAC GGT ACA 201 Pro Gly Val Asn Val Gln Ser Thr Tyr Pro Gly Ser Thr Tyr Ala Ser Leu Asn Gly Thr nydroisse and are especially

661 TCG ATG GCG ACA CCT CAC GTT GCC GGA GCG GCT GCA CTA GTT AAA CAA AAA AAC CCT TCA
useful in detergent formula- 221 Ser Met Ala Thr Pro His Val Ala Gly Ala Ala Ala Leu Val Lys Gla Lys Asn Pro Ser 721 TGG AGC AAC GTT CAA ATC CGC AAC CAC TTA AAA AAC ACA GCT ACT AGC TTA GGC AGT ACT 241 Trp Ser Asn Val Gln Ile Arg Asn His Leu Lys Asn Thr Ala Thr Ser Leu Gly Ser Thr 781 AAC TTA TAC GGC AGC GGC TTA GTT AAC GCT GAA GCT GCA GCT CGT 261 Asn Leu Tyr Gly Ser Gly Leu Val Asn Ala Glu Ala Ala Ara

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-1-

SUBTILISIN MUTANTS

Field of the Invention

The present invention relates to novel carbonyl hydrolase mutants having an amino acid sequence wherein 5 one or more amino acid residues of a precursor carbonyl hydrolase, specifically those at positions corresponding residues +123 and/or +274 in Bacillus amyloliquefaciens subtilisin, have been substituted with a different amino acid. Such mutant carbonyl 10 hydrolases, in general, are obtained by in vitro modification of a precursor DNA sequence encoding a naturally-occurring or recombinant carbonyl hydrolase to encode the substitution of one or both of these amino acid residues in a precursor amino acid sequence alone 15 or in combination with other substitution, insertion or deletion in the precursor amino acid sequence.

Background of the Invention

Serine proteases are a subgroup of carbonyl hydrolase. They comprise a diverse class of enzymes having a wide 20 range of specificities and biological functions. Stroud, R. M. (1974), Sci. Amer., 131, 74-88. Despite their functional diversity, the catalytic machinery of serine proteases has been approached by at least two genetically distinct families of enzymes: the subtilisins and the mammalian chymotrypsin related and homologous bacterial serine proteases (e.g., trypsin and S. gresius trypsin). These two families of serine

proteases show remarkably similar mechanisms of catalysis. Kraut, J. (1977), Ann. Rev. Biochem., 46, 331-358. Furthermore, although the primary structure is unrelated, the tertiary structure of these two enzyme families bring together a conserved catalytic triad of amino acids consisting of serine, histidine and aspartate.

Subtilisin is a serine endoprotease (MW 27,500) which is secreted in large amounts from a wide variety of 10 Bacillus species and other microorganisms. The protein sequence of subtilisin has been determined from at least four different species of Bacillus. Markland, F.S., et al. (1983), Honne-Seyler's Z. Physiol. Chem., 364, 1537-1540. The three-dimensional crystallographic structure 15 of Bacillus amyloliquefaciens subtilisin to 2.5A resolution has also been reported. Wright, C.S., et al. (1969), Nature, 221, 235-242; Drenth, J., et al. (1972), Eur. J. Biochem., 26, 177-181. These studies indicate that although subtilisin is genetically unrelated to the 20 mammalian serine proteases, it has a similar active site structure. The x-ray crystal structures of subtilisin covalently bound peptide inhibitors containing (Robertus, J.D., et al. (1972), Biochemistry, 11, 2439-2449), or product complexes (Robertus, J.D., et al. 25 (1976), <u>J. Biol. Chem.</u>, <u>251</u>, 1097-1103), have also provided information regarding the active site and putative substrate binding cleft of subtilisin. addition, a large number of kinetic and chemical modification studies have been reported for subtilisin 30 (Philipp, M., et al. (1983), Mol. Cell. Biochem., 51, 5-32; Svendsen, B. (1976), Carlsbera Res. Comm., 41, 237-291; Markland, F.S. Id.) as well as at least one report wherein the side chain of methione at residue 222 of subtilisin was converted by hydrogen peroxide to 35 methionine-sulfoxide (Stauffer, D.C., et al. (1965), J. Biol. Chem., 244, 5333-5338) and the side chain of serine at residue 221 converted to cysteine by chemical modification (Polgar, et al. (1981), <u>Biochimica et Biophysica Acta</u>, <u>667</u>, 351-354.)

Patent No. 4,760,025 and EPO Publication 5 No. 0 130 756 published January 9, 1985 each disclose the modification of subtilisin amino acid residues corresponding to positions in Bacillus amyloliquefacien subtilisin tyrosine -1, aspartate +32, asparagine +155, tyrosine +104, methionine +222, glycine +166, histidine 10 +64, glycine +169, phenylalanine +189, serine +33, serine +221, tyrosine +217, glutamate +156 and alanine EPO Publication No. 0 251 446 published +152. January 7, 1988 discloses other amino acid residues in Bacillus amyloliquefaciens subtilisin 15 equivalents which may be modified by way of substitution, insertion or deletion and which may be combined with modifications to the residues identified in U.S. Patent No. 4,760,025 to form useful subtilisin mutants. The particular residues identified herein, 20 however, are not identified in these references.

Similarly, PCT Publication No. W0 89/09819 and W0 89/09830 each published October 19, 1989, disclose subtilisin enzymes made by mutating a nucleotide sequence coding for a subtilisin. Numerous amino acid residues are identified in each of these publications which may be so modified. However, as with the previously identified references, neither identifies the residues of the present invention.

Accordingly, it is an object herein to provide carbonyl hydrolase mutants containing the substitution of amino acid residues in a precursor carbonyl hydrolase corresponding to positions +123 and/or +274 in Bacillus amyloliquefaciens subtilisin. Such mutants generally have at least one property which is different from the

same property of the carbonyl hydrolase precursor from which the amino acid of said mutant is derived.

It is further object to provide DNA sequences encoding such carbonyl hydrolase mutants as well as expression vectors containing such mutant DNA sequences.

still further, another object of the invention is to provide host cells transformed with such vectors as well as host cells which are capable of expressing such DNA to produce carbonyl hydrolase mutants either intracellularly or extracellularly.

Summary of the Invention

The invention includes non-naturally occurring carbonyl hydrolase mutants having a different proteolytic activity, stability, and/or performance characteristic 15 as compared to the precursor carbonyl hydrolase from which the amino acid sequence of the mutant is derived. The precursor carbonyl hydrolase may be a naturallyoccurring carbonyl hydrolase or recombinant hydrolase. Specifically, such carbonyl hydrolase mutants have an amino acid sequence, not found in nature, which is derived by replacement of one or more amino acid residues of a precursor carbonyl hydrolase with one or more different amino acids. The one or more amino acid residues of the precursor enzyme correspond to positions 25 Asn +123 and/or Ala +274 of Bacillus amyloliquefaciens subtilisin or equivalent amino acid residues in other carbonyl hydrolases or subtilisins.

The invention also includes mutant DNA sequences encoding such carbonyl hydrolase or subtilisin mutants.

These mutant DNA sequences are derived from a precursor DNA sequence which encodes a naturally-occurring or recombinant precursor enzyme. The mutant DNA sequences

are derived by modifying the precursor DNA sequence to encode the substitution of one or more specific amino acid residues encoded by the precursor DNA sequence corresponding to position +123 and/or +274 in Bacillus amyloliquefaciens. These recombinant DNA sequences encode carbonyl hydrolase mutants having a novel amino acid sequence and, in general, at least one property which is substantially different from the same property of the enzyme encoded by the precursor carbonyl hydrolase DNA sequence. Such properties include proteolytic activity, stability and/or enhanced performance characteristics.

The invention also includes procaryotic and eucaryotic subtilisins with a different amino acid residue such as serine, at positions equivalent to Asn +123 in <u>Bacillus amyloliquefaciens</u> subtilisin and to subtilisin with different amino acid residues at positions equivalent to position +274 in <u>Bacillus amyloliquefaciens</u> subtilisin.

Further, the invention includes expression vectors containing such mutant carbonyl hydrolase DNA sequences as well as host cells transformed with such vectors which are capable of producing such mutants. The invention also relates to detergent compositions comprising the carbonyl hydrolase mutants of the invention.

Brief Description of the Drawings

Fig. 1 depicts the DNA and amino acid sequence for Bacillus amyloliquefaciens subtilisin and a partial restriction map of this gene.

-6-

Fig. 2 depicts the conserved amino acid residues among subtilisins from <u>Bacillus amyloliquefaciens</u>, <u>Bacillus subtilis</u> varI168 and <u>Bacillus licheniformis</u> (carlsbergensis).

- 5 Figs. 3A and 3B depict the amino acid sequence of subtilisin from <u>Bacillus</u> <u>amyloliquefaciens</u>, <u>Bacillus</u> <u>subtilis</u> varI168 and <u>Bacillus</u> <u>licheniformis</u>.
- Fig. 4 depicts the amino acid sequence of three subtilisins. The top line represents the amino acid sequence of subtilisin from <u>Bacillus amyloliquefaciens</u> subtilisin (also sometimes referred to as subtilisin BPN'). The second line depicts the amino acid sequence of subtilisin from <u>Bacillus lentus</u> (subtilisin 309 in PCT Publication No. WO 89/06276). The bottom line represents the amino acid sequence of a preferred embodiment of the invention designated GG-RYSA. The symbol * denotes the absence of specific amino acid residues as compared to subtilisin BPN'.
 - Fig. 5 depicts the construction of plasmid pGG A274.
- 20 Fig. 6 depicts the construction of pGG-KVNA which is an intermediate to plasmid pGG-RYSA.
 - Fig. 7 depicts the oligonucleotide-duplex method used to construct a synthetic <u>Bacillus lentus</u> subtilisin gene.
- 25 Fig. 8 depicts the strategy for constructing a synthetic gene encoding <u>Bacillus</u> <u>lentus</u> subtilisin.
- Fig. 9 depicts the cassette used to make substitutions in the DNA at codon position +123 by cassette mutagenesis. XXX represents the codon modified to encode the amino acid substitutions at position +123.

Fig. 10 depicts the DNA and amino acid sequence of a preferred embodiment of the invention wherein the DNA sequence is a synthetic DNA. The DNA in this Figure has been modified to encode arginine at position 27, serine at position 78, tyrosine at position 104, serine at position 123 and alanine at position 274.

Detailed Description of the Invention

It has been discovered that in vitro mutations in the carbonyl hydrolase subtilisin at an amino acid residue equivalent to +123 in <u>Bacillus amyloliquefaciens</u> subtilisin produces subtilisin mutants exhibiting altered proteolytic activity over precursor subtilisins.

It has also been discovered that in vitro mutation at residues equivalent to +274 in Bacillus 15 amyloliquefaciens subtilisin produce subtilisin mutants exhibiting altered stability, e.q. modified In some instances, these autoproteolytic stability. latter mutants also exhibit enhanced performance when used in detergent compositions.

20 Carbonyl hydrolases are enzymes which hydrolyze compounds containing

- bonds in which X is oxygen or nitrogen. They include naturally-occurring carbonyl hydrolases and recombinant carbonyl hydrolases. Naturally-occurring carbonyl hydrolases principally include hydrolases, e.g. peptide hydrolases, such as subtilisins or metalloproteases.
 Peptide hydrolases include α-aminoacylpeptide hydrolase, peptidylamino acid hydrolase, acylamino hydrolase,
 - peptidylamino acid hydrolase, acylamino hydrolase, serine carboxypeptidase, metallocarboxypeptidase, thiol proteinase, carboxylproteinase and metalloproteinase.

Serine, metallo, thiol and acid proteases are included, as well as endo and exo-proteases.

"Recombinant carbonyl hydrolase" refers to a carbonyl hydrolase in which the DNA sequence encoding the naturally-occurring carbonyl hydrolase is modified to produce a mutant DNA sequence which encodes the substitution, insertion or deletion of one or more amino acids in the carbonyl hydrolase amino acid sequence. Suitable modification methods are disclosed herein, in EPO Publication No. 0 130 756 published January 9, 1985 and EPO Publication No. 0 251 446 published January 7, 1988.

Subtilisins are bacterial or fungal carbonyl hydrolases which generally act to cleave peptide bonds of proteins As used herein, "subtilisin" means a 15 or peptides. naturally-occurring subtilisin or a recombinant subtilisin. A series of naturally-occurring subtilisins is known to be produced and often secreted by various microbial species. Amino acid sequences of the members 20 of this series are not entirely homologous. However, the subtilisins in this series exhibit the same or similar type of proteolytic activity. This class of serine proteases shares a common amino acid sequence defining a catalytic triad which distinguishes them from 25 the chymotrypsin related class of serine proteases. The subtilisins and chymotrypsin related serine proteases both have a catalytic triad comprising aspartate, histidine and serine. In the subtilisin related proteases the relative order of these amino acids, reading from the amino to carboxy terminus is aspartatehistidine-serine. In the chymotrypsin related proteases the relative order, however is histidine-aspartate-Thus, subtilisin herein refers to a serine protease having the catalytic triad of subtilisin 35 related proteases. Examples include the subtilisins

identified in Fig. 3 herein and as described in PCT Publication W0 89/06276 and EPO Publication No. 0 283 075.

"Recombinant subtilisin" refers to a subtilisin in which
the DNA sequence encoding the subtilisin is modified to
produce a mutant DNA sequence which encodes the
substitution, deletion or insertion of one or more amino
acids in the naturally-occurring subtilisin amino acid
sequence. Suitable methods to produce such modification
and which may be combined with those disclosed herein
include those disclosed in EPO Publication
Nos. 0 130 756 and 0 251 446 and PCT Publication Nos.
W0 89/06279, W0 89/09830 and W0 89/09819.

"Non-human carbonyl hydrolases" and the DNA encoding 15 them may be obtained from many procaryotic and eucaryotic organisms. Suitable examples of procaryotic organisms include gram negative organisms such as $\underline{\mathbf{E}}$. coli or Pseudomonas and gram positive bacteria such as Micrococcus or Bacillus. Examples of eucaryotic 20 organisms from which carbonyl hydrolase and their genes may be obtained include yeast such as Saccaromycees cerevisiae, fungi such as Aspergillus sp., and non-human mammalian sources such as, for example, bovine sp. from which the gene encoding the carbonyl hydrolase chymosin 25 can be obtained. As with subtilisins, a series of carbonyl hydrolases can be obtained from various related species which have amino acid sequences which are not entirely homologous between the members of that series but which nevertheless exhibit the same or similar type 30 of biological activity. Thus, non-human carbonyl hydrolase as used herein has a functional definition refers to carbonyl hydrolases which associated, directly or indirectly, with procaryotic and eucaryotic sources.

A "carbonyl hydrolase mutant" has an amino acid sequence which is derived from the amino acid sequence of a "precursor carbonyl hydrolase". The precursor carbonyl naturally-occurring carbonyl include hydrolases 5 hydrolases and recombinant carbonyl hydrolases. amino acid sequence of the carbonyl hydrolase mutant is "derived" from the precursor hydrolase amino acid sequence by the substitution, deletion or insertion of one or more amino acids of the precursor amino acid 10 sequence. Such modification is of the "precursor DNA sequence" which encodes the amino acid sequence of the precursor carbonyl hydrolase rather than manipulation of the precursor carbonyl hydrolase enzyme per se. Suitable methods for such manipulation of the precursor 15 DNA sequence include methods disclosed herein and in EPO Publication Nos. 0 130 756 and 0 251 446.

Specific residues corresponding to positions +123 and +274 of <u>Bacillus amyloliquefaciens</u> subtilisin are identified herein for substitution. These amino acid position numbers refer to those assigned to the mature <u>Bacillus amyloliquefaciens</u> subtilisin sequence presented in Fig. 1. The invention, however, is not limited to the mutation of this particular subtilisin but extends to precursor carbonyl hydrolases containing amino acid residues at positions which are "equivalent" to the particular identified residues in <u>Bacillus amyloliquefaciens</u> subtilisin.

A residue (amino acid) of a precursor carbonyl hydrolase is equivalent to a residue of <u>Bacillus amyloliquefaciens</u>

30 subtilisin if it is either homologous (i.e., corresponding in position in either primary or tertiary structure) or analogous to a specific residue or portion of that residue in <u>Bacillus amyloliquefaciens</u> subtilisin (i.e., having the same or similar functional capacity to combine, react, or interact chemically).

In order to establish homology to primary structure, the amino acid sequence of a precursor carbonyl hydrolase is directly compared to the Bacillus amyloliquefaciens subtilisin primary sequence and particularly to a set 5 of residues known to be invariant in all subtilisins for which sequence is known (Fig. 2). After aligning the conserved residues, allowing for necessary insertions and deletions in order to maintain alignment (i.e., avoiding the elimination of conserved residues through 10 arbitrary deletion and insertion), the residues equivalent to particular amino acids in the primary sequence of Bacillus amyloliquefaciens subtilisin are Alignment of conserved residues preferably should conserve 100% of such residues. 15 alignment of greater than 75% or as little as 50% of conserved residues is also adequate to define equivalent residues. Conservation of the catalytic triad, Asp32/His64/Ser221 should be maintained.

For example, in Fig. 3 the amino acid sequence of subtilisin from <u>Bacillus amyloliquefaciens</u>, <u>Bacillus subtilis</u> var. I168 and <u>Bacillus lichenformis</u> (carlsbergensis) are aligned to provide the maximum amount of homology between amino acid sequences. A comparison of these sequences shows that there are a number of conserved residues contained in each sequence. These are the residues identified in Fig. 2.

These conserved residues thus may be used to define the corresponding equivalent amino acid residues of <u>Bacillus amyloliquefaciens</u> subtilisin in other carbonyl hydrolases such as subtilisin from <u>Bacillus lentus</u> (PCT Publication No. W089/06279 published July 13, 1989) and the preferred subtilisin mutant herein. These particular amino acid sequences are aligned in Fig. 4 with the sequence of <u>Bacillus amyloliquefaciens</u> subtilisin to produce the maximum homology of conserved

residues. As can be seen there are a number of deletions in the sequence of <u>Bacillus lentus</u> and in the preferred subtilisin mutant of the invention as compared to <u>Bacillus amyloliquefaciens</u> subtilisin. Thus, the equivalent amino acid for Val-165 in <u>Bacillus amyloliquefaciens</u> subtilisin in the other subtilisins is the particular isoleucine shown beneath Val-165.

In Fig. 4, the amino acid at position 123 is asparagine in Bacillus amyloliquefaciens subtilisin. In Bacillus 10 lentus subtilisin the equivalent residue is the particular asparagine shown. In the preferred subtilisin mutant of the invention, however, the amino acid equivalent to +123 in Bacillus amyloliquefaciens subtilisin is an amino acid other than asparagine and 15 is preferably the serine shown in Fig. 4. Similarly, in Fig. 4, the amino acid at position +274 Bacillus amyloliquefaciens subtilisin is alanine. seen, the equivalent amino acid in Bacillus lentus subtilisin is the particular threonine shown in Fig. 4. 20 In a particular preferred subtilisin mutant of the invention, the equivalent amino acid position 274 is occupied by the alanine shown in Fig. 4.

Thus, the positions +123 and +274 are identified by primary amino acid sequences in Fig. 4 for the 25 subtilisin from Bacillus lentus and the preferred embodiment of the invention. However, various other amino acid residues may be modified which are equivalent to specific amino acids in Bacillus amyloliquefaciens subtilisin. Thus, in the preferred embodiment, the 30 amino acid lysine at position 27 in Bacillus amyloliquefaciens subtilisin has an equivalent lysine at position 27 in Bacillus lentus subtilisin. indicated in the Examples, the subtilisin comprising one of the preferred embodiments of the invention was 35 derived by modifying a DNA sequence encoding Bacillus

Such modifications to the DNA lentus subtilisin. included the modification of codons equivalent to positions 123 and 274 of Bacillus amyloliquefaciens subtilisin. However, two other modifications were made 5 to the <u>Bacillus lentus</u> amino acid sequence at positions equivalent to residues 27 and 104 in Bacillus amyloliquefaciens subtilisin. Thus, as can be seen in Fig. 4, the lysine at equivalent residue 27 in Bacillus lentus subtilisin was modified to encode arginine in the 10 preferred embodiment. Similarly, the valine residue at position 104 of Bacillus lentus, which is equivalent to tyrosine 104 in Bacillus amyloliquefaciens subtilisin, was also modified to encode tyrosine. preferred embodiment shown in Fig. 4 contains an amino 15 acid sequence derived from Bacillus lentus subtilisin by modifying residues of that subtilisin equivalent to Bacillus 274 of 123 and 27. 104, positions amyloliquefaciens subtilisin.

Equivalent residues may also be defined by determining 20 homology at the level of tertiary structure for a precursor carbonyl hydrolase whose tertiary structure determined by x-ray crystallography. Equivalent residues are defined as those for which the atomic coordinates of two or more of the main chain 25 atoms of a particular amino acid residue of the hydrolase and Bacillus carbonyl precursor amyloliquefaciens subtilisin (N on N, CA on CA, C on C, and 0 on 0) are within 0.13nm and preferably 0.1nm after alignment. Alignment is achieved after the best model 30 has been oriented and positioned to give the maximum overlap of atomic coordinates of non-hydrogen protein atoms of the carbonyl hydrolase in question to the Bacillus amyloliquefaciens subtilisin. The best model is the crystallographic model giving the lowest R factor 35 for experimental diffraction data at the highest resolution available.

$$R \ factor = \frac{\sum_{h} |Fo(h)| - |Fc(h)|}{\sum_{h} |Fo(h)|}$$

Equivalent residues which are functionally analogous to a specific residue of Bacillus amyloliquefaciens subtilisin are defined as those amino acids of the precursor carbonyl hydrolases which may adopt a 5 conformation such that they either alter, modify or contribute to protein structure, substrate binding or catalysis in a manner defined and attributed to a specific residue of the Bacillus amyloliquefaciens subtilisin. Further, they are those residues of the 10 precursor carbonyl hydrolase (for which a tertiary structure has been obtained by x-ray crystallography), which occupy an analogous position to the extent that although the main chain atoms of the given residue may not satisfy the criteria of equivalence on the basis of 15 occupying a homologous position, the atomic coordinates of at least two of the side chain atoms of the residue lie with 0.13nm of the corresponding side chain atoms Bacillus amyloliquefaciens subtilisin. coordinates of the three dimensional structure of 20 Bacillus amyloliquefaciens subtilisin are set forth in EPO Publication No. 0 251 446 and can be used as outlined above to determine equivalent residues on the level of tertiary structure.

Some of the residues identified for substitution,
insertion or deletion are conserved residues whereas
others are not. In the case of residues which are not
conserved, the replacement of one or more amino acids
is limited to substitutions which produce a mutant which
has an amino acid sequence that does not correspond to
one found in nature. In the case of conserved residues,

such replacements should not result in a naturallyoccurring sequence. The carbonyl hydrolase mutants of
the present invention include the mature forms of
carbonyl hydrolase mutants as well as the pro- and
prepro-forms of such hydrolase mutants. The preproforms are the preferred construction since this
facilitates the expression, secretion and maturation of
the carbonyl hydrolase mutants.

"Prosequence" refers to a sequence of amino acids bound
to the N-terminal portion of the mature form of a
carbonyl hydrolase which when removed results in the
appearance of the "mature" form of the carbonyl
hydrolase. Many proteolytic enzymes are found in nature
as translational proenzyme products and, in the absence
of post-translational processing, are expressed in this
fashion. A preferred prosequence for producing carbonyl
hydrolase mutants, specifically subtilisin mutants, is
the putative prosequence of <u>Bacillus amyloliquefaciens</u>
subtilisin although other subtilisin prosequences may
be used. In the Examples, the putative pro sequence
from the subtilisin from <u>Bacillus lentus</u> (ATCC 21536)
was used.

A "signal sequence" or "presequence" refers to any sequence of amino acids bound to the N-terminal portion of of a carbonyl hydrolase or to the N-terminal portion of a prohydrolase which may participate in the secretion of the mature or pro forms of the hydrolase. This definition of signal sequence is a functional one, meant to include all those amino acid sequences, encoded by the N-terminal portion of the subtilisin gene or other secretable carbonyl hydrolases, which participate in the effectuation of the secretion of subtilisin or other carbonyl hydrolases under native conditions. The present invention utilizes such sequences to effect the secretion of the carbonyl hydrolase mutants as defined

herein. A preferred signal sequence used in the Examples comprises the first seven amino acid residues of the signal sequence from <u>Bacillus subtilis</u> subtilisin fused to the remainder of the signal sequence of the subtilisin from <u>Bacillus lentus</u> (ATCC 21536).

A "prepro" form of a carbonyl hydrolase mutant consists of the mature form of the hydrolase having a prosequence operably linked to the amino-terminus of the hydrolase and a "pre" or "signal" sequence operably linked to the 10 amino terminus of the prosequence.

"Expression vector" refers to a DNA construct containing a DNA sequence which is operably linked to a suitable control sequence capable of effecting the expression of said DNA in a suitable host. Such control sequences 15 include a promoter to effect transcription, an optional operator sequence to control such transcription, a sequence encoding suitable mRNA ribosome binding sites, and sequences which control termination of transcription and translation. The vector may be a plasmid, a phage 20 particle, or simply a potential genomic insert. Once transformed into a suitable host, the vector may replicate and function independently of the host genome, or may, in some instances, integrate into the genome In the present specification, "plasmid" and 25 "vector" are sometimes used interchangeably as the plasmid is the most commonly used form of vector at present. However, the invention is intended to include such other forms of expression vectors which serve equivalent functions and which are, or become, known in 30 the art.

The "host cells" used in the present invention generally are procaryotic or eucaryotic hosts which preferably have been manipulated by the methods disclosed in EPO Publication No. 0 130 756 to render them incapable of

secreting enzymatically active endoprotease. A preferred host cell for expressing subtilisin is the Bacillus strain BG2036 which is deficient in enzymatically active neutral protease and alkaline protease (subtilisin). The construction of strain BG2036 is described in detail in EPO Publication No. 0 130 756 and further described by Yang, M.Y., et al. (1984), J. Bacteriol., 160, 15-21. Other host cells for expressing subtilisin include Bacillus subtilis I168 (EPO Publication No. 0 130 756).

Host cells are transformed or transfected with vectors constructed using recombinant DNA techniques. Such transformed host cells are capable of either replicating vectors encoding the carbonyl hydrolase mutants or expressing the desired carbonyl hydrolase mutant. In the case of vectors which encode the pre or prepro form of the carbonyl hydrolase mutant, such mutants, when expressed, are typically secreted from the host cell into the host cell medium.

"Operably linked" when describing the relationship between two DNA regions simply means that they are functionally related to each other. For example, a presequence is operably linked to a peptide if it functions as a signal sequence, participating in the secretion of the mature form of the protein most probably involving cleavage of the signal sequence. A promoter is operably linked to a coding sequence if it controls the transcription of the sequence; a ribosome binding site is operably linked to a coding sequence if it is positioned so as to permit translation.

The genes encoding the naturally-occurring precursor carbonyl hydrolase may be obtained in accord with the general methods described in EPO Publication Nos. 0 130 756 and 0 251 446. As can be seen from the

examples disclosed therein, the methods generally comprise synthesizing labelled probes having putative sequences encoding regions of the hydrolase of interest, preparing genomic libraries from organisms expressing the hydrolase, and screening the libraries for the gene of interest by hybridization to the probes. Positively hybridizing clones are then mapped and sequenced.

The cloned carbonyl hydrolase is then used to transform a host cell in order to express the hydrolase. 10 hydrolase gene is then ligated into a high copy number plasmid. This plasmid replicates in hosts in the sense that it contains the well-known elements necessary for plasmid replication: a promoter operably linked to the gene in question (which may be supplied as the gene's 15 own homologous promotor if it is recognized, i.e., transcribed, by the host), a transcription termination and polyadenylation region (necessary for stability of the mRNA transcribed by the host from the hydrolase gene in certain eucaryotic host cells) which is exogenous or 20 is supplied by the endogenous terminator region of the hydrolase gene and, desirably, a selection gene such as an antibiotic resistance gene that enables continuous cultural maintenance of plasmid-infected host cells by growth in antibiotic-containing media. High copy number 25 plasmids also contain an origin of replication for the host, thereby enabling large numbers of plasmids to be the cytoplasm without chromosomal limitations. However, it is within the scope herein to integrate multiple copies of the hydrolase gene into 30 host genome. This is facilitated by procaryotic and eucaryotic organisms which are particularly susceptible to homologous recombination.

Alternatively, a synthetic gene encoding a naturallyoccurring or mutant precursor carbonyl hydrolase may be
produced. In such an approach, the DNA and/or amino

acid sequence of the precursor hydrolase is determined. Multiple, overlapping synthetic single-stranded DNA fragments are thereafter synthesized which upon hybridization and ligation produce a synthetic DNA 5 encoding the precursor hydrolase. This approach provides several advantages over cloning the natural gene in that restriction sites may be interposed throughout the DNA without change in the amino acid sequence encoded so as to facilitate subsequent 10 modification to form mutant carbonyl hydrolases. Further, the synthetic approach allows for adjusting the codon usage in the synthetic gene to conform with the codon bias for the particular expression hosts to be used. An example of synthetic gene construction is set 15 forth in the Examples.

Once the naturally-occurring or synthetic precursor carbonyl hydrolase gene has been cloned, a number of modifications are undertaken to enhance the use of the gene beyond synthesis of the naturally-occurring precursor carbonyl hydrolase. Such modifications include the production of recombinant carbonyl hydrolases as disclosed in EPO Publication Nos. 0 130 756 and 0 251 446 and the production of carbonyl hydrolase mutants described herein.

25 The following cassette mutagenesis method may be used to facilitate the construction and identification of the carbonyl hydrolase mutants of the present invention although other methods including site-directed mutagenesis may be used. First, the naturally-occurring 30 gene encoding the hydrolase is obtained and sequenced in whole or in part. Then the sequence is scanned for a point at which it is desired to make a mutation (deletion, insertion or substitution) of one or more amino acids in the encoded enzyme. The sequences 35 flanking this point are evaluated for the presence of

restriction sites for replacing a short segment of the gene with an oligonucleotide pool which when expressed will encode various mutants. Such restriction sites are preferably unique sites within the hydrolase gene so as 5 to facilitate the replacement of the gene segment. However, any convenient restriction site which is not overly redundant in the hydrolase gene may be used, provided the gene fragments generated by restriction digestion can be reassembled in proper sequence. 10 restriction sites are not present at locations within a convenient distance from the selected point (from 10 to 15 nucleotides), such sites are generated by substituting nucleotides in the gene in such a fashion that neither the reading frame nor the amino acids 15 encoded are changed in the final construction. Mutation of the gene in order to change its sequence to conform to the desired sequence is accomplished by M13 primer extension in accord with generally known methods. task of locating suitable flanking regions 20 evaluating the needed changes to arrive at convenient restriction site sequences is made routine by the redundancy of the genetic code, a restriction enzyme map of the gene and the large number of different restriction enzymes. Note that if a convenient flanking 25 restriction site is available, the above method need be used only in connection with the flanking region which does not contain a site.

Once the naturally-occurring DNA or synthetic DNA is cloned, the restriction sites flanking the positions to be mutated are digested with the cognate restriction enzymes and a plurality of end termini-complementary oligonucleotide cassettes are ligated into the gene. The mutagenesis is enormously simplified by this method because all of the oligonucleotides can be synthesized so as to have the same restriction sites, and no

synthetic linkers are necessary to create the restriction sites.

As used herein, proteolytic activity is defined as the rate of hydrolysis of peptide bonds per milligram of active enzyme. Many well known procedures exist for measuring proteolytic activity (K. M. Kalisz, "Microbial Proteinases", Advances in Biochemical Engineering/Biotechnology, A. Fiechter ed., 1988).

In one aspect of the invention, the objective is to 10 secure a mutant carbonyl hydrolase having a greater (numerically large) proteolytic activity as compared to the precursor carbonyl hydrolase, thereby enabling the use of the enzyme to more efficiently act on a target substrate. Specific amino acids useful to obtain such 15 results in subtilisin-type carbonyl hydrolases at residues equivalent to +123 in Bacillus amyloliquefaciens subtilisin are presented in the Examples. In some instances, lower proteolytic activity may be desirable. In such cases a decrease in 20 proteolytic activity can be produced by substituting the amino acids identified in the examples at residues equivalent to +123 in Bacillus amyloliquefaciens subtilisin.

For precursor subtilisins wherein serine is not the 25 residue at the position equivalent to +123 in Bacillus amyloliquefaciens the greatest proteolytic activity is obtained when serine is substituted in the precursor at position +123. Further, no naturally-occurring Bacillus subtilisin is known to exist which contains serine at 30 a position equivalent to +123 in Bacillus amyloliquefaciens subtilisin. Based on the discovery that serine at this position enhances proteolytic activity, one skilled in the art can screen naturallyoccurring Bacillus subtilisin to identify and clone a

natural mutant containing serine at this position. Such natural <u>Bacillus</u> subtilisin mutants are within the scope of the invention.

Where the carbonyl hydrolase is from other than <u>Bacillus</u>
and a serine is present at +123 in the precursor enzyme
the substitution can be one that decreases proteolytic
activity. This would be useful, for example, where the
synthetic activity of the carbonyl hydrolases is desired
(as for synthesizing peptides). One may wish to
decrease this proteolytic activity which is capable of
destroying the product of such synthesis.

In another aspect of the invention, it has been determined that residues equivalent to +274 in Bacillus amyloliquefaciens subtilisin are important in modulating 15 the overall performance characteristics of the enzyme in detergent compositions. Thus, as set forth in the Examples, the threonine in Bacillus lentus subtilisin at equivalent position +274 can be mutated to alanine in the preferred embodiment to produce enhanced 20 performance of the mutant enzyme. As also disclosed in the Examples, substitution of this residue with an amino acid other than threonine, e.g. leucine, serine, valine and alanine results in a decrease in the stability of the mutant. Such decrease in stability is believed to 25 be the result of autocatalytic degradation of the mutant. Thus, modifications of residues equivalent to +274 in Bacillus subtilisin are capable of enhancing the overall performance of the enzyme in a detergent composition and modulating the overall stability of the 30 enzyme. In this aspect of the invention, the objective is to secure a mutant carbonyl hydrolase having enhanced performance when used in a detergent composition as compared to the precursor carbonyl hydrolase. As used herein, enhanced performance in a detergent is defined 35 as increased cleaning of certain enzyme sensitive stains

such as grass or blood. This cleaning is determined by visual evaluation after a standard wash cycle.

A preferred embodiment of the invention is set forth in the Examples wherein the lysine at position 27 is substituted with arginine, the valine at position 104 is substituted with tyrosine, the asparagine at position 123 substituted with serine and the threonine at residue 274 is substituted with alanine in Bacillus lentus subtilisin. Although the stability of this enzyme is somewhat reduced as compared to the precursor Bacillus lentus subtilisin, the performance level of this enzyme in a detergent composition is substantially enhanced such that the same performance of this Bacillus lentus subtilisin mutant is obtained as compared to the unmodified Bacillus lentus subtilisin when using approximately one-half the amount of enzyme.

Based on the results obtained with this and other mutant subtilisins, it is apparent that residues in carbonyl hydrolases equivalent to positions +123 and +274 in Bacillus amyloliquefaciens are important to the proteolytic activity, performance and/or stability of these enzymes.

Many of the carbonyl hydrolase mutants of the invention, especially subtilisin, are useful in formulating various detergent compositions. A number of known compounds are suitable surfactants useful in compositions comprising the carbonyl hydrolase mutants of the invention. These include nonionic, anionic, cationic, anionic, or zwitterionic detergents, as disclosed in U.S. 4,404,128 to Barry J. Anderson and U.S. 4,261,868 to Jiri Flora, et al. The art is familiar with the different formulations which can be used as cleaning compositions.

Subtilisins of the invention can be formulated into known powdered and liquid detergents having pH between 6.5 and 12.0 at levels of about .01 to about 5% preferably .1% to .05%) by weight. These detergent cleaning compositions can also include other enzymes such as known proteases and amylases, as well as builders and stabilizers.

The addition of subtilisins of the invention to conventional cleaning compositions does not create any special use limitation. In other words, any temperature and pH suitable for the detergent is also suitable for the present compositions as long as the pH is within the above range, and the temperature is below the subtilisins of the invention denaturing temperature. In addition, subtilisins of the invention can be used in a cleaning composition without detergents, again either alone or in combination with builders and stabilizers.

The following is presented by way of example and is not to be construed as a limitation to the scope of the claims.

EXAMPLE 1

Constructions for Expression of Bacillus lentus
Subtilisin Gene in Bacillus subtilis

Plasmid pSAR, Fig. 5, carries a translational fusion via a common Sau3A restriction site at the seventh/eighth signal sequence codon of the subtilisin genes of B. subtilis and B. amyloliquefaciens. As shown in Fig. 5, this gene, on an EcoRI-BamHI 2.0 Kb fragment, was subcloned into M13mp19 in order to isolate single-stranded template DNA to be used for site-directed mutagenesis to form pSAR-Q275R. The mutagenesis protocol was essentially that of Zoller, M., et al.

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(1983), <u>Methods Enzymol.</u>, <u>100</u>, 468-500, (1) and used a synthetic oligonucleotide of the sequence:

5'-C-AAC-GTA-CAG-G<u>CT-GCA-G</u>CT-CGC-TAA-AAC-ATA-A-3'
O275R

where the asterisks denote changes from the wild-type gene sequences and the underline represents an introduced PstI restriction endonuclease site used in screening for the particular mutant gene encoding the Q275R change. These changes were made to (1) convert the amino acid at this position to that found in Bacillus lentus subtilisin and (2) to allow hookup of the terminator in pSAR to the mature coding region of Bacillus lentus via a Pst site similarly introduced into pGG36 from Bacillus lentus (ATCC 21536).

Plasmid pGG36, Fig. 5, contains a 2.1 kb genomic DNA fragment from <u>Bacillus lentus</u> (ATCC 21536) encoding the complete subtilisin gene which was cloned by standard methods in the shuttle vector pBS42. Band, L., et al. (1984), <u>DNA</u>, <u>3</u>, 17-21.

The amino acid sequence for this subtilisin is the same as that disclosed for subtilisin 309 in PCT Publication No. 89/06279. This gene was subcloned into M13 as above for site-directed mutagenesis using an oligonucleotide of the sequence:

5'-C-AAT-GCA-GAA-G<u>CT-GCA-G</u>CT-CGC-TAA-TCA-A-3' T274A

in order to 1) introduce a PstI site at the same 30 location in this gene corresponding to the site introduced into pSAR above and 2) to substitute the threonine at position 274 with alanine to form pGG36-T274A.

The mutant pSAR-Q275R and pGG36-T274A genes were individually subcloned back into pBS42 prior to PstI/BamHI digestions, fragment isolation and ligation to produce plasmid GG-A274B.amy.term. as shown in Fig. 5, all by standard methods.

A synthetic DNA linker was made by annealing complimentary single-stranded oligonucleotides of the sequences:

5'-G-ATC-GTC-GCG-TCG-ACC-GCA-CTA-CTC-ATT-TCT-GTT-GCT-TTT-AGT-TCA-T-3'

and

10

5'-CGA-TGA-ACT-AAA-AGC-AAC-AGA-AAT-GAG-TAG-TGC-GGT-CGA-CGC-GAC-3'

to give the double-stranded DNA fragment #2 shown in Fig. 6. The recessed left- and right-hand ends of this duplex linker are complimentary to the Sau3A end of fragment #1 (from pSAR) and the ClaI end of fragment #3 (from pGG-A274 B.amy.term), respectively. These 3 fragments were combined with fragment 4 from pSAR-Q275R after restriction endonuclease digestions of plasmids, fragment isolation and ligation by standard methods to produce plasmid pGG-KVNA. The designation GG-KVNA indicates that this subtilisin contains the subtilisin encoded by pGG-36 which includes lysine (K) at position 27, valine (V) at position 104, asparagine (N) at position 123 and the substitution of threonine at position 274 with alanine (A).

EXAMPLE 2

Modification of PGG-KVNA

As indicated in Fig. 6, the GG-KVNA gene (2.1 kb EcoRI-BamHI fragment) was subcloned into M13 for three successive rounds of site-directed mutagenesis using oligonucleotides having the sequence:

- (a) 5'-GT-TCT-GGT-GTA-AGA-GTT-GCT-GT<u>T-CTA-GA</u>T-ACA-GGT-3',
 K27R
- (b) 5'-A-GTA-TTA-GGG-<u>GCT-AGC</u>-GGT-TCA-GGT-TCG-TAC-AGC-TCG-ATT-3' V104y
 and
- 15 (c) 5'-GGG-AAC-AAT-GGA-ATG-CAC-GTT-GCT-AGC-TTG-AGT-TTA-3'

The asterisks denote changes from the wild-type gene sequence. The underlines represent, in (a) an introduced XbaI site and in (b) and (c) introduced NheI sites used to screen for the presence of the linked R27, Y104 and S123 mutations, respectively. In addition, in (c), the overlined denotes a destroyed SphI site. Finally, the 2.1 kb GG-RYSA gene was subcloned back into pBS42 for expression in B. subtilis hosts.

The resultant plasmid was designated pGG-RYSA. This designation indicates that four residues were modified in the pGG-KVNA plasmid. Lysine (K) at position 27 to arginine (R), valine (V) to tyrosine (Y) at position 104 and asparagine (N) at position 123 to serine (S). The alanine previously substituted at residue 274 was not modified in this procedure.

The lysine at position 27 was substituted with arginine based upon the amino acid sequencing of subtilisin 309. As indicated in PCT Publication No. W089/06279, lysine is located at position 27. However, after independently 5 sequencing this subtilisin protein, the initial data indicated that arginine was the residue at position 27. In the case of the substitution of tyrosine for valine at residue 104, the substitution was made to lower the pH activity profile and to increase the performance of 10 the enzyme based on results previously obtained for subtilisin Bacillus amvloliquefaciens referred to as BPN'). The substitution of asparagine at position 123 with serine is based on the results obtained hereinafter wherein it was determined that the 15 substitution of serine at position 123 maximized the proteolytic activity of the enzyme in a closely related mutant.

EXAMPLE 3

Construction of Synthetic Bacillus lentus Subtilisin Gene

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DNA encoding the amino acid sequence of <u>Bacillus lentus</u> subtilisin was also prepared by constructing a gene encoding a synthetic DNA sequence.

The 2.1 kb HindIII genomic fragment from plasmid pGG36
was sequenced. The deduced amino acid sequence of the
mature gene product (GG36 subtilisin) was used to design
a synthetic mature coding sequence with the following
properties: (1) In general, the codons most frequently
found for each amino acid in seven different <u>B. subtilis</u>
genes (from a tabulation of codon usages, Table 2 from
Maruyama, T., et al., (1986), <u>Nucl. Acids Res.</u>,
Supplement <u>14 pp. r151-r197</u>) were utilized except in the
cases where alternate codons resulted in conveniently

located restriction enzyme recognition sites within the gene; (2) Approximately every 40-60 b.p. of the ~0.8 mature coding region, combinations of 2 or 3 specifically chosen codons were utilized which resulted 5 in the introduction of fairly evenly spaced, unique restriction sites. These sites were chosen to facilitate (a) later cassette mutagenesis and screening studies and (b) constructions involving more than one mutation; (3) A unique Pst I recognition site was 10 designed to cover codons 272-274 allowing hook up to the terminator sequences of a <u>Bacillus</u> <u>amyloliquefaciens</u> gene similarly modified over the same three codons and substituting threonine at position 274 with alanine; and (4) A unique NruI site was introduced to cover mature 15 codons residues 9-10 allowing hookup to GG36's pre-pro coding sequence via a short synthetic duplex DNA linker. Based on this design, oligonucleotides ("oligos") were synthesized such that upon annealing the coding and noncoding oligos for a given -60 b.p. coding region, the 20 resultant duplex DNA fragment would have at it's ends single stranded regions complimentary to the end of the next duplex fragment of the gene (see, Fig. 7).

A total of 36 separate oligos (comprising 18 individual duplexes) were used in the scheme, as outlined above, resulting in an -0.8kb duplex synthetic mature coding region (Fragment 3 in Fig. 8).

Finally, one additional pair or synthetic oligo's was synthesized, which upon annealing (to give fragment 2 in Fig. 8) has an NcoI site at it's 5' end (complimentary to GG36's NcoI site at mature codons 5-6) and an NruI site at its 3' end (complimentary to the 3's 5' end of fragment 3).

The final construction to give a complete expression unit consisting of <u>B. subtilis</u> promoter and the first

seven amino acids of the signal sequence hooked up to GG36's sequences encoding the remainder of the signal sequence, the complete pro sequence and the first six mature amino acids (Fragment 1 from GG-KVNA), the synthetic gene encoding mature residues 7-274 (Fragments 2+3) and the terminator region (including the final mature gene codon 279) of Bacillus amyloliquefaciens (fragment 4) was done as a four-way ligation as set forth in Fig. 6.

10 Finally, three additional separate mutations were introduced into the mature coding region of this full length hybrid gene. The first substituted the lysine at position 27 with arginine. The second substituted the valine at position 104 with tyrosine. The third substituted the asparagine at position 123 with serine The resultant plasmid is designated pBC3-RYSA. The following example describes the method used to modify position 123 in the synthetic gene. Similar methods were used to modify positions 27 and 104 in this synthetic gene.

EXAMPLE 4

Construction of Position 123 Mutants

An Xho I site was introduced over codons 111/112 in the synthetic gene from Example 3 by making three phenotypically silent mutations via site directed mutagenesis (primer extension mutagenesis in M13). The resulting plasmid, pX123 (Fig. 9), was digested with Xho I and Ava I and the large vector-containing fragment isolated by electroelution from agarose gel.

Complimentary synthetic oligonucleotides were annealed, ligated with the pX123 large fragment and transformed into E. coli strain MM294. These cassettes encoded, individually, all 20 naturally-occurring amino acids at

position 123, and in addition contained a silent mutation which destroyed a unique Sph I site lying between the Xho I and Ava I sites in pX123. Resulting plasmids from E. coli transformants were screened for the loss of the unique Sph I site. Positives by restriction analysis (i.e., Sph I negatives) were sequenced to confirm the presence of the desired position 123 mutations subcloned into the shuttle vector pBS42 and transformed into Bacillus subtilis BG2036 for expression.

EXAMPLE 5

Activity of Various +123 Mutants

Proteolytic activity of each of the subtilisin mutants encoded by the above modified position +123 mutants was assayed by mixing 0.04 ml of supernatant from centrifuged culture broths with 0.56 ml of 1% w/v casein in 0.1M Tris pH8.60. After a 20 minute incubation at 37°C, reactions were quenched by precipitation with 10% trichloroacetic acid (TCA). Activity was determined from the absorbance at a wavelength of 280nm for the supernatant after precipitation with 10% TCA.

TABLE I

Relative proteolytic activity of codon 123

variants normalized to Asn-123 mutant

5	Codon 123	<pre>% Proteolytic Activity</pre>
	Ser	116
	Asn	100
	Cys	22
	Gly	12
10	Ala	9
	Thr	7
	Gln	7
	Val	6
	Glu	<5
15	Ile	<5
	Trp	<5
	Phe	<5
	Asp	<5
	His	<5
20	Leu	<5
	Met	<5
	Pro	<5
	Tyr	<5

In the process of final confirmation of the DNA sequence of the synthetic gene coding for the enzyme BC3-RYSA, proline was found to be at position 78 instead of serine (the amino acid at this position in <u>Bacillus lentus</u> subtilisin). The initial properties of the position 123 mutations were tested in this enzyme, BC3-RPYA (proline at position 78). These results are shown in Table I. The amino acid at position 78 was thereafter changed back to serine to form the DNA and amino acid sequence shown in Fig. 10 by replacing the synthetic DNA duplex corresponding to that portion of the gene.

As can be seen the substitution of Asn with Ser at position +123 results in a substantial increase in proteolytic activity. The relationship between the various subtilisins discussed herein are summarized for positions 27, 78, 104, 123 and 274 in Table II.

TABLE II position

Ī		27	78	104	123	274
Ì	GC36 (genomic)	Lys(K)	Ser(S)	Val(V)	Asn(N)	Thr(T)
10	Synthetic B. lentus gene	Lys(K)	Pro(P)	Val(V)	Asn(N)	Ala(A)
	B.amylolique- facins subtilisin (BPN)	Lys(K)	Ser(S)	Tyr(Y)	Asn(N)	Ala(A)
15	Subtilisin 309 as published	Lys(K)	Ser(S)	Val(V)	Asn(N)	Thr(T)
	Preferred embodiment herein	Arg(R)	Ser(S)	Tyr(Y)	Ser(S)	Ala(A)

EXAMPLE 6

Stability of Position 274 Mutants

20 Stability of position 274 mutants in BC3-RPY (arginine at position 27, proline at position 78, and tyrosine at position 104 in <u>Bacillus lentus</u> subtilisin) are shown in Table III. Data are percent activity remaining following incubation at 37°C in 50mM EDTA for 60 minutes.

TABLE III

	Amino Acid at Position 274	% Activity
	Leucine	2%
5	Serine	79%
	Threonine	91%
	Valine	42%
	Alanine	43%

Mutations at this position clearly effect stability of
the enzyme. Although the alanine mutation was not as
stable as serine or threonine at this position, this
enzyme provided superior performance relative to
Bacillus lentus subtilisin under the conditions of use
described. For different applications, other amino
acids at position 274 may be used.

EXAMPLE 7

<u>Detergent Composition</u>

A spray-dried phosphate detergent granule of the following composition was prepared:

	Component	Weight &
	Sodium C12 linear alkylbenzene sulfonate	8.45
	Sodium Tallow Alcohol sulfate	4.23
5	Sodium C14~15 linear alkyl sulfate	4.23
	Sodium Toluene Sulfonate	1.00
	Sodium Tripolyphosphate	5.60
	Sodium pyrophosphate	22.40
	Silicate (1.6 r)	5.50
10	Sodium Sulfate	29.83
	Sodium polyacrylate (4500 MW)	1.17
	Brightener	0.22
	Sodium Carbonate	12.30
	Polyethylene Glycol (MW 8000)	0.47
15	C12 ⁻ 13 alcohol polyethoxylate (6.5)*	0.50
	Miscellaneous + Water	to 100%
	Protease**	0.034

*Alcohol and monoethoxylate alcohol removed.

20 **mg active enzyme/g (2.0 mg active enzyme/g stock)

A 0.1 weight percent solution of this composition in water had a pH of 10.0. The composition with subtilisin mutant of the invention (Fig. 7) provided superior cleaning of enzyme-sensitive stains, when compared to Bacillus lentus at 0.068 mg active enzyme/g product, in a 95°F (35°C) wash at 6 grains per gallon (gpg) hardness (3:1 Ca/Mg).

Throughout this application reference is made to various amino acids by way of common one-and three-letter codes.

30 Such codes are identified in <u>Proteins: Structures and Molecular Proteases</u>, Thomas E. Creighton, eds. W.N. Freeman, N.Y., N.Y. (1983), p.3.

Although the preferred form of the invention has been described above, it will be obvious to those skilled in the art to which the invention pertains, that, after understanding the invention as a whole, various changes and equivalent modifications may be made without departing from the scope of the invention as defined by the appended claims.

All publications are expressly incorporated herein by reference.

WHAT IS CLAIMED IS:

- A carbonyl hydrolase mutant having an amino acid sequence not found in nature which is derived form a precursor carbonyl hydrolase by substituting a different amino acid for the amino acid residue at a position in said precursor equivalent to +123 or +274 in <u>Bacillus amyloliquefaciens</u> subtilisin.
 - 2. The carbonyl hydrolase mutant of Claim 1 wherein said precursor carbonyl hydrolase is a subtilisin.
- 10 3. The subtilisin mutant according to Claim 2 wherein said substitution is at a position equivalent to +123.
- 4. The subtilisin mutant of Claim 3 wherein the amino acid residue in said subtilisin mutant at said 15 position is serine.
 - 5. The subtilisin mutant according to Claim 2 which is derived from a <u>Bacillus</u> subtilisin.
- 6. A mutant <u>Bacillus</u> subtilisin exhibiting improved proteolytic activity which is derived from naturally20 occurring or mutant precursor <u>Bacillus</u> subtilisin which has the amino acid residue at a position equivalent to +123 in <u>Bacillus</u> amyloliquefaciens subtilisin changed to serine.
- 7. A <u>Bacillus</u> subtilisin having serine at a position 25 equivalent to +123 in <u>Bacillus</u> <u>amyloliquefaciens</u> subtilisin.

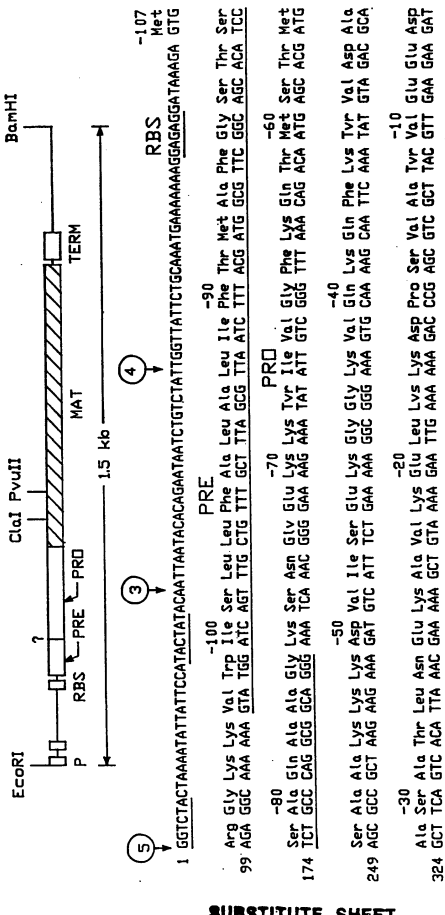
8. A mutant <u>Bacillus</u> subtilisin having the amino acid sequence:

AQSVPWGOSRVQAPAAHNRGLTGSGVRVAVLDTGISTHPDLNIRGGASFVPGE
PSTQDGNGHGTHVAGTIAALNNSIGVLGVAPSAELYAVKVLGASGSGSYSSIA
QGLEWAGNNGMHVASLSLGSPSPSATLEQAVNSATSRGVLVVAASGNSGAGSI
SYPARYANAMAVGATDQNNNRASFSQYGAGLDIVAPGVNVQSTYPGSTYASLN
GTSMATPHVAGAAALVKQKNPSWSNVQIRNHLKNTATSLGSTNLYGSGLVNAE
AAAR.

- 9. DNA encoding the carbonyl hydrolase mutant of 10 claim 1.
 - 10. Expression vector encoding the DNA of Claim 9.
 - 11. Host cell transformed with the expression vector of Claim 10.
- 12. An enzymatic cleaning composition capable of degrading proteins comprising:
 - a) a surfactant end; and
 - b) a carbonyl hydrolase mutant having an amino acid sequence not found in nature which is derived form a precursor carbonyl hydrolase by substituting a different amino acid for the amino acid residue at a position in said precursor equivalent to +123 or +274 in <u>Bacillus</u> amyloliquefaciens subtilisin.
 - 13. The composition of Claim 12 wherein said carbonyl hydrolase enzyme comprises a subtilisin.
- 25 14. The composition of Claim 13 wherein said subtilisin has the following amino acid sequence:

AQSVPWGOSRVQAPAAHNRGLTGSGVRVAVLDTGISTHPDLNIRGGASFVPGE PSTQDGNGHGTHVAGTIAALNNSIGVLGVAPSAELYAVKVLGASGSGSYSSIA OGLEWAGNNGMHVASLSLGSPSPSATLEQAVNSATSRGVLVVAASGNSGAGSI SYPARYANAMAVGATDQNNNRASFSQYGAGLDIVAPGVNVQSTYPGSTYASLN GTSMATPHVAGAAALVKQKNPSWSNVQIRNHLKNTATSLGSTNLYGSGLVNAE AAAR.

- 15. The composition of Claim 12 wherein the 5 surfactant comprises a detergent.
 - 16. A composition according to Claim 15 comprising a spray-dried detergent granule.



Ala. GCC Gtn CAA Va l GTA Ser TCT His Leu TTA HIS LEu CTG Asn GAT Thr Ala GCT 45 CCT G1< Pro CCT His HIS Ala Ser TCT Ser TCT Lys AAA Asn AAC Ser TCT 10 Gln 11e L CAA ATT A Asp GAT Asp Asn AAC 60 Asp GAC Ile Val Ser GTA TCA **Gly GGT** Gla Ser AGC Phe TTC 61y 660 Asp GAC Asn Pro CCT 30 Val Ile (GTT ATC (Pro Asn AAT Tyr Pro CCT Thr ACA ۷a (GTG Ala GCG Gtu GAA Ser TCC Val GTA Ser TCT Gln CAG Pro CCT Lys AAA -1 | 1 Tyr Ala (TAC GCG (Val GTT Val GTT Asn 50 Met ATG Ala GCG Ser TCA Ser AGC HIS 9Ly 66A Ala GCC Ala GCA Thr ACT G1y GGA Ba l GTA Tyr TAC GI AGC 399

Lvs AAA Va l GTA Ala GCT Tvr TAC 90 Leu CTT Ala Ser TCA Ser Ala GCA Ser AGC Pro CCA Ala GCG Val GTT 61y 660 Val Leu GTA TTA 80 GLy GGT I le ATC Ser TCA Asn Asn AAT Leu CTT Ala GCT Val ACA ACA 70 613 660 624

Met ATG Asn AAT Asn AAC I le ATC Ala GCG Trp TGG Glu GAG 11e ATC 110 Gly GGA Asn AAC 1 le ATT Ile ATC Trp 156 Ser AGC Tyr TAC CA G 61y 660 Ser TCC 100 Gly GGT Ala Asp GAC Asp Ala GCT **G**17 Leu CTC Va l GTT 669

Ala GCA Va l GTT Ala GCC Lvs AAA 140 Asp GAT Val GTT Ala GCA A la GCG Lvs AAA Leu Ala GCT Ala GCT Ser TCT **G**1y 130 Ser TCT Pro CCT G(y 61y 660 Leu CTC Ser AGC Met ATG Asn AAC I le ATT 120 Asp GAC 774

5 6 6 Pro CCT TVr TAC Va t GTG Thr ACA Ser AGC Ser TCA Ser AGC 160 Gly GGC Thr Ser TCC Ser Thr ACT 61y 660 Asn Glu 61y 66T Ata Ata GCA Ala GCG 150 Val GTT Val GTC Va l GTA Val GTC Ser TCC

549

				U	
Pro	G (<	Thr	Asn AAC	icTo	
G1 v GGA	Asn	Asn AAC	I le ATC	דככנ	
Vsl GTA	TVL	Thr	Leu CTG	TCAA	
Ser AGC	Ala Tvr / GCG TAC /	Trp TGG	G (v	ATGT	
190 Ser TCA	61y 666	240 Asn AAC	Lvs	ວຍວວ	
Phe TTC	Tvr TAC	r S	G1 v GGA	тсст	
Ser TCT	Lvs AAA	His	Tyr	11CT	
A La GCA	Asn AAC	Lvs AAG	Tyr	ATTT	
Arp AGA	G1v GGA	Ser	Phe Tyr Tyr TTG TAC TAT	TATT	
Gla Arp	210 Pro CCT	Leu Ile Leu Ser Lvs TTG ATT CTT TCT AAG	260 Ser TCT	TERM C <u>ataaaaaaccgcc</u> cttggccccgccgcttttttttttcttcctccgcatgttcaatccgctcc	
Asn AAC	Leu	Ile ATT	\sp iAT	C66T	
Ser Asn AGC AAC	Thr	Leu TTG	1 61y A	7 500 160	
Ser	Ser AGC	Ala GCT	Leu	TERM	
Asp GAC	Gln	Ala GCT	Lvs	T T	
180 Val GTT	I le ATC	230 Aa l GCG	Thr		
Ala GCT	Ser TCT	Gly GGA	Thr	AAAA	
G (\	Va l GTA	Ala GCC	Thr	ATAA	
Val	פנא פפכ	Va l GTT	Asn AAC	AAC	
Ala	Pro CCT	His	GAA GAA	DC TAA	
11e ATT	200 Ala GCA	Pro CCG	250 Leu TTA	275 Gln CAG	
Val GTC	Met ATG	Ser TCT	Ser AGT	Ala GCT	
Ser TCT	Va l GTC	Ala GCA	Ser	Ala GCA	
Pro CCT	Asp GAT	Met ATG	Aro CGC	Ala GCG	
TVT	Leu Asp CTT GAT	Ser Met Ala TCA ATG GCA	Val Aro GTC CGC	Gln CAA	
170 Lvs AAA	Glu	220 Thr ACG	Gln	270 Val GTA	
924	666	1074	1149	1224	
				. 4	

1416 CTTCCCGGTTTCCGGTCAGCTCAATGCCATAACGGTCGGCGGCGTTTTCCTGATACCGGGAGACGGCATTCGTAATCGGATC

1316 ATAATCGACGGATGGCTCCCTCTGAAAATTTTAACGAGAAACGGCGGGTTGACCCGGCTCAGTCCCGTAACGGCCAACTCCTGAAACGTCTCAATCGCCG

FIG.-1C

MARTITUTE CUPPY

TD'	TAL	LY	CON	NSE!	RVE	D F	RES	IDU	ES]	IN	SUE	TI	LIS	ZNI	•				20
•	•	•	•	Р	•	•	•	•		•	•		•	•			•		
21	•	G	•	1		•	•		30	•	D	•	G	•			•	Н	40
41			•	•	G	•	•	•	50 ·	٧		•	•	,	•	•	•	•	60
61	•	•	Н	G	Т	Н	•		70 G			•		•			•		80
81		G		•	•	•	1	•	90	•	•	•	•	٧	L			•	100 G
1 0 1 S		•			•		•		110 G		•			•	•	•	•		120
121	•	•			L	G		•	130		•	•	•	•		•			140
141	•	•	•	•	G	ı	•		150			•	G	N	•	•	•	•	160
161	•	ı		•		٧	P	•	170			•				٧	•	•	180
181	•	•	•	•	•		S	F	190 S		•	•	•		•	•	•	•	500
201 P	G	ı		•	•	•	•	•	210		•	•			•	•		G	220 T
221 S	М	Α	•	Р	Н	٧	Α	G	530		•	•	•	•	•		•	•	240
241		•	•	•	•	R	•		250		•	•	•	•		•	•	•	260
261 ·	•		•					N	270 •		•	•	•	•					

FIG.-2

Homology of Bacllur proteases

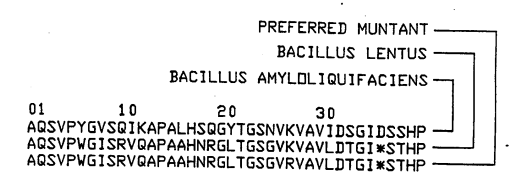
- 1.Bacillus amyloliquifaciens 2.Bacillus subtilis var.I168 3.Bacillus licheniformis (carlsbergensis)

		•	_									_							
1 A A	Q Q Q	2 2 T	V V V	P P P	Y Y Y	G G	V I I	\$ \$ P	10 Q Q L	I I I	K K K	A A A	P P D	A A K	L L V	H H Q	S S A	Q Q Q	20 G G
21 Y Y F	T T K	G G	2 2 A	N N N	V V V	K K K	V V V	A A A	A A A 30	I I L	D D D	S S T	G G	1 1 1	D D Q	2 2 A	S S S	H H H	40 P P P
41 D D D	L L L	K N N	V V V	A R V	G G G .	G G	A A A	2 2	50 M F F	V V V	P P A	S G	EEE	T T A	N N Y	P P N	F Y T	Q Q	60 D D D
61 N D G	N L N	S N G	H V H	· G R G	T G T	H G H	V A V	A S A	70 G F G	T V T	V P V	A S A	A E A	L T L	N N D	N P N	S Y T	I Q T	80 G D G
81 V V V	L L L	G G	V V V	A S A	P P P	2 2	A A V	S S	9(L L L) Y Y Y	A A A	V V V	K K K	V V V	L L L	G D N	A 2 2	D T S	100 G G G
1 (2 2 3	01 G G	Q Q S	Y Y Y	2 2 2	W W G	1 1 1	I I V	N S	1 : G G	1 0 1 1	E E E	VV	A A A	I I T	A S T	N N N	N N G	М М М	120 D D D

FIG.-3A

12 V V V	I I I	N N N	M M M	2 2	L L L	G G	G G	P P A	13 S T S	0 G G	S S S	A T T	A A A	L L M	K K K	A T Q	A V A	V V V	140 D D D
14 K K N	1 A A A	V V Y	A S A	S S R	G G	V I V	V V V	V V V	15 V A V	0 A A	A A A	A A A	G G	N N N	E S	G G	T S N	2 2	160 G G
16 S S S	1 S T T	S 2 N	T T	V V I	G G	Y Y Y	P P P	G A A	17 K K K	0 Y Y	P P D	2 2	V T V	I I I	A A A	V V V	G G	A A A	180 V V V
18 D N D	1 S S	S	. 2 N	Q Q N	R R R	A A A	2 2 2	F F F	19 S A S	0 2 2 3	G A V	v G	P S A	E E E	L L	D D E	V V V	M M M	200 A A A
20 P P P	1 G G	V V A	S V	I I G	Q Q Y	2 2	T T T	L L Y	21 P P P	0 G G T	N G N	K T T	Y Y Y	G G A	A A T	Y Y L	N N N	G G	220 T T T
2 2 2 5 55	1 M M M	2 A A	A T S	P P P	H H	V V V	A A A	G G	23 A A	80 A A	A A A	L L L	I I I	L L L	2 2	K K K	H H N	P P P	240 N T N
24 W W L	1 T T S	N N A	T A S	Q Q Q	V V V	R R R	S D N	S R R	25 L L	50 E S	N S S	T T	T A A	T T T	K Y Y	L L L	G G	Д И 2	2 2 2 560
26 F F F	1	Y Y Y	G G	K K K	G G	L L I	I	N N N	27 V V V	0 Q Q E	A A A	A A. A	A A A	Q Q Q					

FIG. -3B substitute sheet



41 50 60 70
DLKVAGGASMVPSETNPFQDNNSHGTHVAGTVAALNNSIG
DLNIRGGASFVPGE*PSTQDGNGHGTHVAGTIAALNNSIG
DLNIRGGASFVPGE*PSTQDGNGHGTHVAGTIAALNNSIG

81 90 100 110
VLGVAPSASLYAVKVLGADGSGQYSVIINGIEWAIANNMD
VLGVAPSAELYAVKVLGASGSGSVSSIAQGLEWAGNNGMH
VLGVAPSAELYAVKVLGASGSGSYSSIAQGLEWAGNNGMH

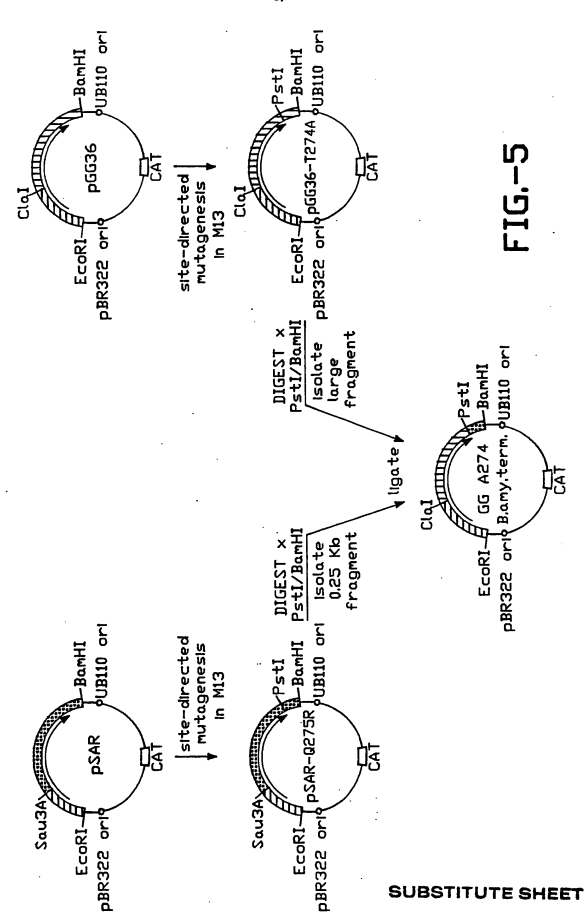
121 130 140 150
VINMSLGGPSGSSAALKAAVDKAVSGVVVVAAAGNEGTSG
VANLSLGSPSPSATLEQAVNSATSRGVLVVAASGNSGAGS
VASLSLGSPSPSATLEQAVNSATSRGVLVVAASGNSGAGS

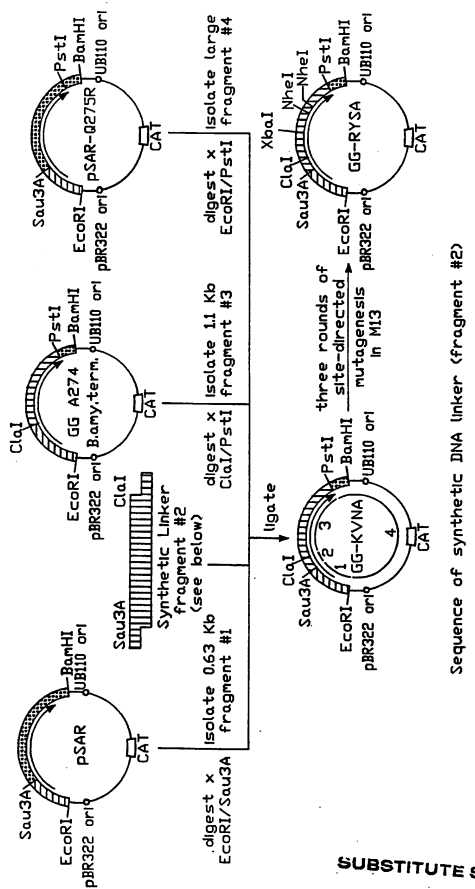
161 170 180 190 SSSTVGYPGKYPSVIAVGAVDSSNQRASFSSVGPELDVMA ****ISYPARYANAMAVGATDQNNNRASFSQYGAGLDIVA ****ISYPARYANAMAVGATDQNNNRASFSQYGAGLDIVA

201 210 220 230
PGVSIQSTLPGNKYGAYNGTSMASPHVAGAAALILSKHPN
PGVNVQSTYPGSTYASLNGTSMATPHVAGAAALVKQKNPS
PGVNVQSTYPGSTYASLNGTSMATPHVAGAAALVKQKNPS

241 250 260 270
WTNTQVRSSLENTITKLGDSFYYGKGLINVQAAAQ
WSNVQIRNHLKNTATSLGSTNLYGSGLVNAEAAAR
WSNVQIRNHLKNTATSLGSTNLYGSGLVNAEAAAR

FIG. -4 SUBSTITUTE SHEET





5'-G-ATC-GTC-GCG-TCG-ACC-GCA-CTA-CTC-ATT-TCT-GTT-GCT-TTT-AGT-TCA-T-3' 3'-CAG-CGC-AGC-TGG-CGT-GAT-GAG-TAA-AGA-CGA-AAA-TCA-AGT-AGC-5'

SUBSTITUTE SHEET

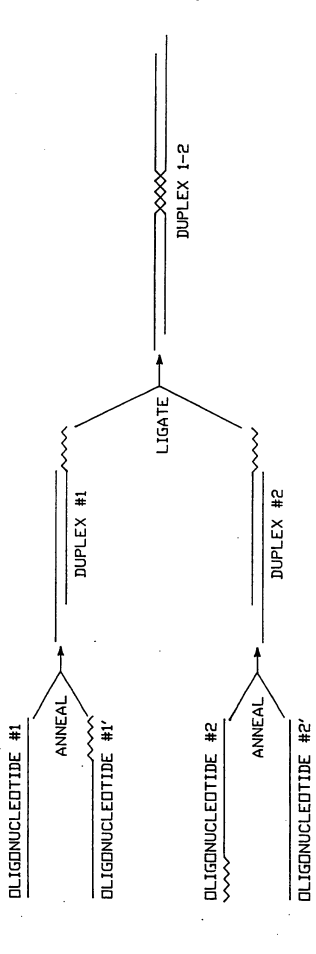
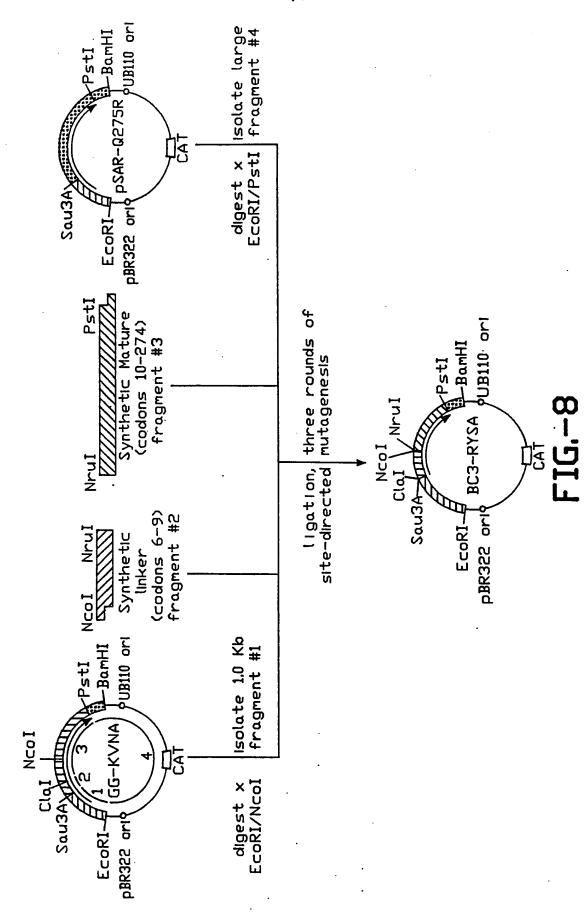


FIG.-7



>	AGC CTC GGG-3' TCG GAG CCC-5' AvaI	CTC GGG-3' GAG CCC-5 Ava!	pTC GGG-3; C-5′	XXX TTA AGC CTC GGG-3' XXX AAT TCG GAG CCC-5' 123
פֿ	C GGC G CCC Ava I	, GG , CC , a	99 ;	ָטָט טפּ
Leu	CTC GAG	CTC GAG Av	ptc	CTC
Ser	AGC TCG	AGC TCG		AGC TCG
Leu	TTA AAT	TTA AAT		TTA AAT
113 Ala Gly Asn Asn Gly Met His Val Ala Asn Leu Ser Leu Gly	F AAC TTA	AAC TTA AGC TTG AAT TCG	•	XXX 123
Ala	GCT CGA	GCT		GCT
Val	GGC ATG CAC GTT GCT CCG TAC GTG CAA CGA	AAC AAC GGC ATG CAC GTT GCT TTG TTG CCG TAC GTG CAA CGA Sph1		* GGC AAC AAC GGT ATG CAC GTT GCT CCG TTG TTG CCA TAC GTG CAA CGA SphI destroyed
120 HIS	CAC	CAC		CAC GTG stroy
Me.	ATG TAC	ATG TAC		ATG TAC des
Gly	000 000 000	200 200 200 200		GGT CCA Sph1
Asn	AAC TTG	AAC TTG		AAC TTG
Asn	AAC AAC	AAC TTG		AAC TTG
115 61y	099	999 090		900 000
Ala	GCT	GCT CGA		GCT CGA
Trp	TGG ACC	TGG ACC	_	
פות	GAA	* * * CTC GAG GAG CTC XhoI	СТР	C GAG G CTC XhoI
Leu	TTA GAA AAT CTT	CTC GGAG CXAOI	C GAG CTP	CTC GAG ★
110 Gly Leu Glu Trp	993 993	922	900 999	GGC CTC GAG TGG CCG GAG CTC ACC Xhol
- •	<u>-</u> ∢	- «	ut I .	213 */ DNA es:
W.T. A.A.	W.T. DNA	pX123 DNA	pX123 cu w/ XhoI and AvaI	Cut.pX1213 [1gated w/ duplex DNA cassettes:
7.	۲. ۲	pX18	pX123 cut w/ XhoI and AvaI	Cut Liga dup cass

FIG.-9

66C 61y

CGC

AAC Asn

CAC

GCT

GCT Ala

CCT

GCT

CAA

GTT Val

CGA

TCG Ser

ATC Ite

66C 61y

TGG Trp

CCA

GTG Val

TCA

CAA

CCA

CAC

١

ACA

AGC Ser

ATC Ile

66C 61y

ACA. Thr

GAT Asp

TTA

GTT Val

GCT Ala

GTT

AGA

GTT

66C 61y

AGC Ser

66C 61y

ACA

TTA

61 21 GAT Asp

ACA

TCG

CCG

GAA

66C 61y

CCC Pro

GTT Val

AGC Ser

666 Ala

66C 61y

66C 61y

AGA Arg

ATT Ile

AAT Asn

CTT

GAT Asp

121 41 66A 61y

ATC Ite

TCG

AAC Asn

AAC

TTA

GCT

GCT

ATC Ile

ACA

66C 61y

GCC Ata

GTT Val

CAC H1s

ACA

66C

CAC His

66C 61y

AAC Asn

66C 61y

181 .61 66C 61y

AGC Ser

GCT Ala

66C 61y

TTA

GTT Va(

AAA Lys

GTT Val

GCT Ala

TAC

TTA

GAA

GCC Ala

TCG Ser

CCT

GCT

GTT Val

66C 61y

TTA

GTT Val

241 81 CAC

AGT Met

667 61y

AAC Asn

AAC Asn

66C 61y

GCT Ala

766 Trp

GAG Glu

CTC Leu

66C 61y

CAA Gln

GCT Ala

ATC Ile

TCT

AGC

TAC

AGC Ser

66C 61y

AGC Ser

301

361 121

GTT Val

GCT

CAA

GAA TTA ACA GCT AGC Ser CCT Pro AGC Ser CCT Pro AGC Ser 666 61y CTC AGC Ser TTA AGC Ser GCT Ala GTT Val

TCG Ser	ACA Thr		ACA Thr	TCA Ser	ACT Thr	
GGA G l _' y	CGC Ala	GTG Vα (66T 6 t y		AGT Ser	
GCT	766	ATC	AAC	AAC	66C	
Ala	G l y	Ile	Asn	Asn	61y	
66C 61y	767 Va (TTG Leu			٠
AGC Ser	66C A ta	TTA	AGC	CAA Glu		
AAC Asn		66C 61y	GCC	AAA Lys	ACT Thr	CGT Arg
66C	TGC	GCT	ACA TAC GCC	GTT	GCT	GCT
61y	Ala	Ala	Thr Tyr Ala	Val		Ala
AGC	TAA	66C	ACA	CTA	ACA	GCA
	Asn	6 ty	Thr	Leu	Thr	Ala
GCG	CGC	TAC	AGC	GCA	AAC	GCT
Ata	Ala		Ser	A la	Asn	Ala
GCT Ala	ATA Tyr	CAA	CCT GGC Pro Gly	GCT Ala	AAA Lys	GAA
GGC GTT TTA GTT GTT GCT Gly Val Leu Val Val Ala	TAG Arg	AGT	CCT Pro	GCG Ala	TTA	GCT Ala
GTT Val	TGC Ala	TTC	TAC	66A 61y	CAC H1s	AAC Asn
TTA	CCC Pro	AGC Ser	CAA AGC ACA Gin Ser Thr	CAC GTT GCC GGA GCG HIS Val Ala Gly Ala	AAC Asn	Val
GTT	CTA	GCA	AGC	GT 7	ATC CGC	TTA
Val		Ala	Ser	Vα l	Ile Arg	Leu
66C	AGC	AGA	CAA	CAC	ATC	66C
6 t y		Arg	Głn	H1s	Ile	6 ty
AGA	ATC	AAC	GTT	CCT	CAA	AGC
Arg	Ile	Asn	Val		Gln	Ser
TCT	1	AAC	AAC	ACA	GTT	66C
Ser		Asn	Asn	Thr	Val	61y
ACA	1	AAC	GTT	GCG	AAC	TAC
Thr		Asn	Va (Ala	Asn	Tyr
GCT Ala	1	CAA Gln	660 61y	ATG Met	AGC Ser	TTA
AGC	1	GAT	CCT	TCG	766	AAC
Ser		Asp	Pro	Ser	Trp	Asn
421	481	541	601	661	721	781
141	161	181	201		241	261
		•				

FIG.-10B

INTERNATIONAL SEARCH REPORT

International Application No

PCT/US90/06084

. CLASSI	FICATION OF SUBJECT MATTER (if several classifi-	International Application No PCI/US9U/U0U84
	to International Patent Classification (IPC) or to both Natio	
	: C12N 9/48; CO7H 15/12; CO7 K	
	435/212; 536/27; 530/350; 252/8	
	SEARCHED	7.1
		aton Country of
Class-Gastia	Minimum Document	
Classificatio	n System C	lassification Symbols
***	435/91 172.3, 212, 252.3	1. 832:
US.	536/27; 530/350; 252/89.	
	330/2/, 330/330, 232/03.	1
	Documentation Searched other the to the Extent that such Documents	ian Minimum Documentation Bre Included in the Fields Searched ⁵
	CUENTOAT ADORDAORD DAMA DAGE CO	40) 1047 1001
	CHEMICAL ABSTRACTS DATA BASE (C.	
	KEY WORDS: BACILLUS; SUBTILISM	N; CARBONYL HYDROLASE, MUTANT
	MENTS CONSIDERED TO BE RELEVANT 14	
ategory •	Citation of Document, 16 with Indication, where appr	
1		LL et al) 26 July 1988,
Y	see abstract and column	a 3, line 12 to column 1-16
- 1	4, line 55.	2 10
- 1	• • • • • • • • • • • • • • • • • • •	
Y.P	US, A. 4,914,031 (ZUKO)	SSKI et al) 03 April 1-16
1,1	1990, see abstract and	
- 1		
1	and lines 36-45 and co	lumn 3, lines 7-34.
Y	EP, A, 0,130,756 (BOTT	et al), 09 January $1-16$
1	1985, see abstract.	
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* Special	categories of cited documents: 15	"T" later document published after the international filling dat
"A" doci	ument defining the general state of the art which is not	or priority date and not in conflict with the application be cited to understand the principle or theory underlying th
	sidered to be of particular relevance	invention
	er document but published on or after the international grate	"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to
"L" doci	ument which may throw doubts on priority claim(s) or	involve an inventive step
	th is cited to establish the publication date of another ion or other special reason (as specified)	"Y" document of particular relevance; the claimed invention
"O" docu	ument referring to an oral disclosure, use, exhibition or	cannot be considered to involve an inventive step when the document is combined with one or more other such document.
othe	r means	ments, such combination being obvious to a person skille in the art.
"P" docu	Iment published prior to the international filing date but than the priority date claimed	"4" document member of the same patent family
	FICATION	
	Actual Completion of the International Search 2	Date of Mailing of this International Search Report 8
Aera oi rue	- Actom Completion of the International Search *	
		20 FE 1881
	15 January 1001	· AUC LANGE
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